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Thanks in advance,

Peter Brunovskis Art Unit 1632 CM1-Rm 12E05 305-2471

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Involvement of mouse *Mlh1* in DNA mismatch repair and meiotic crossing over

Sean M. Baker^{1*}, Annemieke W. Plug^{2*}, Tomas A. Prolla³, C. Eric Bronner¹, Allie C. Harris¹, Xiang Yao⁴, Donna-Marie Christie¹, Craig Monell⁵, Norm Arnheim⁴, Allan Bradley³, Terry Ashley² & R. Michael Liskay¹

Mice that are deficient in either the *Pms2* or *Msh2* DNA mismatch repair genes have microsatellite instability and a predisposition to tumours. Interestingly, *Pms2*-deficient males display sterility associated with abnormal chromosome pairing in meiosis. Here mice deficient in another mismatch repair gene, *Mlh1*, possess not only microsatellite instability but are also infertile (both males and females). *Mlh1*-deficient spermatocytes exhibit high levels of prematurely separated chromosomes and arrest in first division meiosis. We also show that Mlh1 appears to localize to sites of crossing over on meiotic chromosomes. Together these findings suggest that *Mlh1* is involved in DNA mismatch repair and meiotic crossing over.

¹Department of Molecular and Medical Genetics, L103, Oregon Health Sciences University, 3181 S. W. Sam Jackson Park Road, Portland, Oregon 97201-3098, ŬSA ²Department of Genetics, Yale University School of Medicine, 333 Cedar Street, New Haven, Connecticut 06510, USA ³Department of Molecular and Human Genetics, Howard Hughes Medical Institute, Baylor College of Medicine, Houston, Texas 77030, USA ⁴Molecular Biology Program, University of Southern California, Los Angeles, California 90089-1340, USA ⁵PharMingen, 10975 Torreyana Rd. San Diego, California 92121, USA

*S.M.B. & A.W.P. contributed equally to this study

Correspondence should be addressed to R.M.L. e-mail: liskaym@ ohsu.edu DNA mismatch repair plays a prominent role in the correction of replicative mismatches which escape DNA polymerase proof-reading, mismatches that arise due to spontaneous deamination of 5-methylcytosine, and mispairs that form during genetic recombination. Three genes, *mutS*, *mutL* and *mutH* are central to the correction of replication errors in *Escherichia coli*. Biochemical studies indicate that the MutS protein recognizes DNA mismatches and that the MutH protein is an endonuclease that helps direct repair to newly synthesized strands^{1,2}. The MutL protein appears to couple mismatch recognition by MutS to MutH activation².

Multiple genes with roles in DNA mismatch repair have been identified in the yeast Saccharomyces cerevisiae and in mammals³⁻¹⁰. Genetic analysis in yeast has identified three mutS homologues, MSH2 (ref. 3), MSH3 (ref. 11) and MSH6 (ref. 12), and two mutL homologues, PMS1 and MLH1 (refs 4,5,13,14) that appear to function as components of the same replica-

tive DNA mismatch repair pathway. Recent studies suggest that early steps in yeast mismatch repair include recognition of mispaired bases by a heterodimer of MSH2 and either MSH3, or MSH6 (ref. 12) followed by binding of a heterodimer of MLH1 and PMS1 (ref. 15). In humans, mutation in the *mutS* homologue, *MSH2* (refs 7,8), or in any of three *mutL* homologues, *PMS2*, *PMS1* and *MLH1* (refs 6,9,10), is associated with

hereditary colorectal cancer. Mice engineered to be deficient in either *Pms2* (the homologue of yeast *PMS1*) or *Msh2* show microsatellite instability and predisposition to early onset cancers, primarily lymphomas^{16–18}. The conservation of eukaryotic DNA mismatch repair mechanisms is suggested by biochemical studies that indicate the human MSH2 protein can bind mispaired DNA¹⁹, and that human MLH1 and PMS2 can exist as a heterodimer (ref. 20; A.B. Buermeyer & R.M.L., unpublished observations).

In addition to reducing mutations, DNA mismatch repair appears to play several roles in genetic recombination^{21–24}, including the correction of heteroduplex DNA²¹ and the regulation of recombination between similar but non identical DNA sequences^{25–28}. Moreover, specific DNA mismatch repair gene homologues are involved in different aspects of recombination. For example, the yeast *mutS* homologues, *MSH4* and *MSH5*, each play a part in meiotic crossing over but not DNA mismatch repair^{29,30}. In contrast, yeast *PMS1*

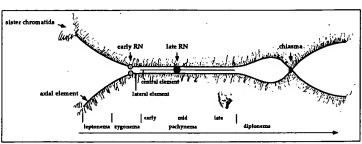
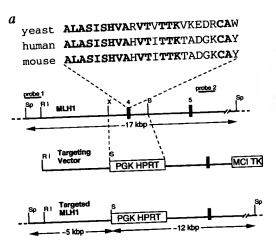


Fig. 1 Meiotic prophase in mammals, showing a timeline of meiotic-specific structural components.



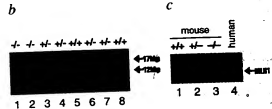


Fig. 2 Generation of Mlh1-deficient mice. a, Scheme used to target the Mlh1 locus in embryonic stem cells. The targeting vector contains approximately 9 kb of homology with the endogenous Mlh1 locus and following integration replaces a 2.5-kb fragment containing exon 4 with the HPRT mini-gene. Expanded above exon 4 is the portion of the Mlh1 protein encoded by this exon. A comparison with mouse, human and S. cerevisiær-Mlh1 proteins indicates that this portion is highly conserved. Bold letters indicate positions of identity in all three proteins. b, Hybridization analysis on DNA from offspring of Mlh1^{+/-} mice. DNA samples were digested with Spel and Sall and genotyped with probe 2. Animals homozygous for the Mlh1 Δ103–126 mutation are pre-

sent in lanes 1 and 2. $Mlh1^{+/+}$ (lanes 5, 8) and $Mlh1^{+/-}$ (lanes 3, 4, 6, 7) animals were also identified. The genotypes have been confirmed by PCR (see Methods). c, Immunoblot analysis on extracts from mouse $Mlh1^{+/+}$, $Mlh1^{+/-}$ and $Mlh1^{+/-}$ fibroblasts cells. Mlh1 protein can be detected in the $Mlh1^{+/+}$ (lane 1) and $Mlh1^{+/-}$ (lane 2) cell extracts, but not in extracts from $Mlh1^{-/-}$ cells (lane 3).

is involved in DNA mismatch repair but not meiotic crossing over¹⁴. Therefore, mismatch repair deficiency in mammals might affect meiotic events such as crossing over, or chromosome pairing. In fact, *Pms2*-deficiency in the mouse results in male infertility associated with the disruption of normal chromosome synapsis (pairing) during meiosis¹⁶.

In most eukaryotes, synapsis and crossing over of homologous chromosomes are critical for the basic function of meiosis: to segregate a complete haploid set of chromosomes to each gamete. Proper segregation of homologues in first division meiosis requires that they are held together by chiasmata to ensure correct chromosomal alignment at metaphase I31,32. Chiasmata represent the physical manifestations of crossing over, or reciprocal recombination, and are first detected cytologically when homologues begin to separate at diplonema (Fig. 1). However, studies on diverse organisms suggest that crossing over occurs earlier in prophase, during pachynema, within electron dense structures on the synaptonemal complex (SC) termed late recombination nodules (late RNs). The number and distribution of late RNs correlate well with crossover events in microorganisms, insects and plants³³ but not in mammals³⁴. Although RNs are assumed to represent recombination protein complexes, specific components of RNs have not been identified.

To define the role of *Mlh1* in mice we have taken two complementary approaches. First, we have derived mice defective in *Mlh1* function and second, we have localized Mlh1 protein in male and female meiotic nuclei. Here we report that in addition to compromising replication fidelity, *Mlh1*-deficiency appears to cause both male and female sterility associated with reduced levels of chiasmata. Significantly, we find that Mlh1 protein appears to be present at sites of interhomologue crossing over and chiasma formation in normal spermatocytes and oocytes. Our findings suggest roles for mouse *Mlh1* in both DNA mismatch repair and meiotic crossing over.

Generation and analysis of Mlh1-deficient mice We used a portion of the human MLH1 cDNA6 as a

probe to isolate mouse *Mlh1* cDNA and genomic clones. To generate mice with a null mutation in *Mlh1*, we transfected a replacement vector (Fig. 2a) into mouse embryonic stem (ES) cells to delete by homologous recombination an exon encoding a highly conserved region (Fig. 2a) of the mouse Mlh1 protein. Targeted ES cell clones were identified by hybridization with probes 5' and 3' (Fig. 2a) to the targeted region (data not shown), and injected into host blastocysts to generate chimaeric animals that transmitted the *Mlh1* mutant allele (Δ103–126 mutation) to F1 offspring (Fig. 2b). Of 145 offspring from *Mlh1*+/- intercrosses, 10% were *Mlh1*+/+, 65% *Mlh1*+/- and 25% *Mlh1*-/-, suggesting that *Mlh1* function is not essential for normal somatic development.

To determine the consequence of the Δ103-126 mutation on Mlh1 protein expression, we used a monoclonal antibody raised against human MLH1 for immunoblot analysis of Mlh1+/+, Mlh1+/- and Mlh1-/fibroblast cells extracts. Consistent with a null mutation, we detected no normal-length Mlh1 in the Mlh1-/- cell extract (Fig. 2c). To determine the consequence of Mlh1 disruption on mismatch repair, we measured mutation in (CA)_n microsatellite repeats using single molecule dilution PCR on spleen DNA samples from Mlh1+/+ and Mlh1-/- animals³⁵. Mlh1deficiency resulted in an elevated level of CA-repeat mutation in spleen DNA (Table 1). Studies on germ cell enriched samples taken from seminiferous tubules demonstrated that Mlh1-deficiency also destabilized simple sequence repeats in the male germ line (Table 1). The elevated mutation level in enriched spermatocytes may represent either mitotic instability, meiotic instability, or both. Preliminary experiments with cultured cell lines from Mlh1-deficient mice show an increased rate of mutation to oubain resistance, suggesting that the role for Mlh1 in replicative fidelity is not limited to microsatellite sequences (C.E.B. & R.M.L., unpublished observation). Furthermore, consistent with the cancer redisposition seen in *Pms2*and Msh2-knockout mice16-18, an Mlh1-deficient female, sacrificed at six months of age for ovarian analysis, had developed a lymphoma.

Table 1 Microsatellite mutation in normal and Mlh1-deficient mice

DNA	Locus	
	D9Mit67	D1Mit355
Mlh1+/+ Spleen	0/103 (0%)	3/131 (2.3%)
Mlh1-/- Spleen	42/221 (19%)	50/246 (20.3%)
^a Mlh1+/+ Sperm	0/136 (0%)	not tested
Mlh1-/- Spermatocytes	32/229 (14%)	33/180 (18.3%)

The number of DNA samples with altered microsatellite sequences divided by the total number of samples examined for each locus. Purified DNA was diluted so that one third of the samples gave a PCR product. The data were obtained through analysis of a single animal of each genotype. The MIh1-/- animal was heterozygous for the C57BL/6 and 129 alleles at each locus. At D9Mit67, we estimate that the alleles differ by 6 (CA/GT) repeats and at D1Mit355 by 7 repeats. In the MIh1-/- mouse, length alterations were observed equally among the C57BL/6 and 129 alleles. At each locus we observed approximately twice as many contractions as expansions, primarily one or two repeats. ^aWe reported this data for wild-type sperm previously16.

Gonadal analysis of Mlh1-deficient animals

To determine the effect of Mlh1-deficiency on the male germline, we examined the contents of the caudal epididymis and testes from $Mlh1^{+/-}$ and $Mlh1^{-/-}$ animals. Whereas the epididymis from Mlh1+/- animals contained normal spermatozoa, we observed no spermatozoa in three Mlh1^{-/-} animals. Histological sectioning of testes from Mlh1-/- animals revealed abnormal spermatogenesis, the most notable phenotype being an absence of late stage cells—round spermatids excess of primary spermatocytes, many with condensed chromosomes, consistent with meiosis I arrest to desynapse, although they remain held together at

in late pachynema, metaphase, or both. Although not directly tested, the total absence of spermatozoa in the Mlh1-/- mice predicts that Mlh1-deficiency result in male infertility.

Matings of Mlh1+/- females with Mlh1+/- males produced litters of normal size (data not shown). However er, matings involving Mlh1-/- females were unsuccessful, suggesting that these females were infertile. Histological sections showed that ovaries from Mlh1-deficient adult females (Fig. 3d) were clearly smaller than wild-type (Fig. 3c), had very few follicle and showed an increased proportion of stromal cells Nonetheless, we observed a reduced number of corpora lutea suggesting that ovulation can occur in the Mlh1-deficient females, albeit infrequently.

A meiotic defect in Mlh1-deficient males

The excess of primary spermatocytes and lack of spermatids in the Mlh1-deficient males suggested a meiosis I arrest. Therefore, we examined meiotic progression in Mlh1-deficient spermatocytes using both standard DAPI staining and immunostaining with Rad51 antibody³⁶. One of the several positive reactions of this polyclonal antibody is the staining of axial (unsynapsed)/lateral (synapsed) elements of the SCs. This reaction, along with the two types of Rad51 foci, provides cytological landmarks for meiotic prophase from leptonema, prior to homologue synapsis, through axial disassembly as homologues begin to separate at diakinesis (see Fig. 1). Rad51-staining revealed that chromosome pairing behavior in the zygotene through early pachytene stages of prophase I was normal in Mlh1-deficient spermatocytes (data not shown). We did not observe the synaptic abnormalities reportedthrough spermatozoa (Fig. 3a,b). There was a clear for Pms2-deficient spermatocytes16. By mid-pachynema in normal mice, the X and Y chromosomes begin

> their distal ends, presumably by a chiasma^{26,37-41} in the psuedoautosomal region (Fig. 4a). In contrast, by mid-pachynema in Mlh1-deficient nuclei, the X and Y were completely separated, giving rise to X and Y univalents (Fig. 4b). The autosomes in both wild-type and Mlh1-/- nuclei remained fully synapsed throughout pachynema (Fig. 4a, b). As homologues desynapse in normal diplonema, the bivalents remained connected at the sites of chiasmata (Fig. 4c). In contrast, as desynapsis of the autosomes progressed in Mlh1-/diplotene nuclei, we observed a steady increase in the frequency of univalents (Fig. 4d). By metaphase I, homologous autosomes of wildtype mice remained connected by chiasmata (Fig. 4e), whereas in Mlh1-deficient nuclei we observed primaris univalents, or unassociated homologues (Fig. 4f). Examination of multiple diplotene and metaphase nuclei indicated that chiasma formation in Mlh1-defi-

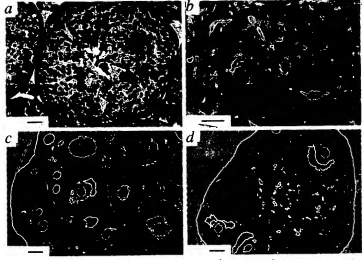
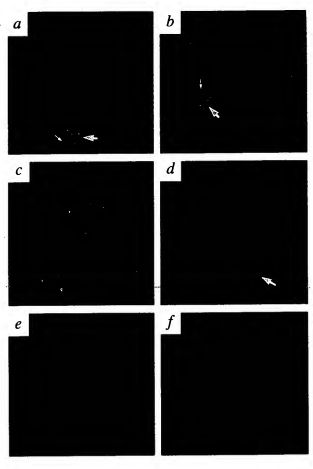


Fig. 3 Testicular and ovarian histology from Mlh1+/- and Mlh1-/- mice. Cross section through $Mlh1^{+/-}$ (a) and $Mlh1^{-/-}$ (b) seminiferous tubules, stained with periodic acid, Schiff's and haematoxylin. Spermatogenesis is normal in the $Mlh1^{+/-}$ tubules (a) as indicated by the production of mature spermatozoa and the appropriate proportion of cells in each sub-stage. In MIh1-1- tubules (b), there is an accumulation of primary spermatocytes and an absence of late stage cells, round spermatids through spermatozoa. The later stages of spermatogenesis are absent from all tubules. Bar in (a) and (b) is 25 μm in length. Histology of Mlh1+/- (c) and Mlh1-/- (d) ovaries stained with eosin and haematoxylin. Note the smaller ovary size and the reduced number of follicles in the mutant (a) as compared to the $Mlh1^{+/-}$ ovary (c). Bar in (c) and (d) is 100 μ m in



cient spermatocytes is reduced 10- to 100-fold. Taken together, these observations suggest that *Mlh1* is required for normal levels of chiasma formation, or stabilization, or both during meiosis I.

Localization of MIh1 during meiosis

We immunostained surface spread spermatocytes and oocytes-from normal mice with an Mlh1 monoclonal antibody and observed discreet fluorescent foci in

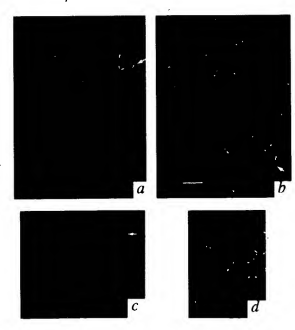


Fig. 4. Analysis of primary spermatocytes from Mih1+ and Mih1+ mice. Spermatocytes were stained with human Rad51 antibody (panels a-d). Left panels represent normal mice, right panels, Mih1+ mice. a,b, Pachytene. a, Wild-type nucleus: the X and Y (arrow) have desynapsed, but show end association; b, Mih1+ nucleus: the X (large arrow) and Y (small arrow) chromosomes are completely separated. Synapsis of the autosomes for wild-type and Mih1-deficient nuclei is completely normal. c, d, Diplotene. c, In wild-type nuclei the autosomes show initial desynapsis, and are held together as bivalents (pairs of homologues) by chiasma. d, In Mih1+ nuclei homologue separation results in univalents (large arrows). e, f, Metaphase stained with DAPI. Brightly staining areas are centromeric heterochromatin. The wild-type nuclei (e) contains 20 bivalents; whereas Mih1+ nuclei (f) contain mostly univalents.

meiotic prophase nuclei. We observed no Mlh1 foci at any stage of meiotic prophase in spermatocytes from Mlh1-/- males indicating that the foci are dependent on Mlh1 and suggesting specificity of the antibody. To provide a temporal and spatial framework for the Mlh1 immunoreactions during meiosis, we used differentially labelled and detected polyclonal antibody³⁶ directed against human Rad51.

In oocytes from normal females double-stained with the Rad51 and Mlh1 antibodies, we observed Mlh1 foci on synapsed portions of the SC as early as zygonema. By the time the homologues were fully synapsed in early pachynema, there were an average of 65 ± 12 Mlh1 foci per oocyte nucleus (n = 9), or 3.25 foci per SC bivalent (Fig. 5d). (Mlh1 foci are rapidly lost during this substage, hence the high standard deviation.) The number of foci decreased to an average of 31 ± 2 Mlh1 foci per nucleus (n = 21), or 1.5 foci per SC bivalent by mid-pachynema and then stabilized. In spermatocytes, we did not see Mlh1 foci on the SCs until homologues were fully synapsed in early pachynema. The number of Mlh1 foci on the SCs in spermatocytes then increased to average of 22 ± 1.5 foci per nucleus (n = 29), or 1.2 foci per autosomal SC bivalent, in mid-pachynema (Fig. 5a). In early pachytene spermatocytes, we consistently observed an Mlh1 focus at the distal end of the pairing region of the X and Y (Fig. 5a). However, no focus was evident at slightly later substages of pachynema as the X and Y began to desynapse precociously relative to the autosomes (Fig. 5b). The number of Mlh1 foci decreased gradually in both oocytes and spermatocytes in late pachynema. In spermatocytes all Mlh1 foci have disappeared by late pachynema. However, in oocytes some Mlh1 foci remain present at sites of chiasmata into early diplonema (Fig. 5c).

Discussion

Using gene targeting in ES cells we derived mice deficient in the DNA mismatch repair gene homologue, Mlh1. We observed increased mutation in microsatellite DNA sequences in both the spleen and the male

Fig. 5 Double labelling with the polyclonal Rad51 antibody (white) and the monoclonal Mlh1 antibody (red). a, Early pachytene spermatocyte- small arrows show Mlh1 foci on SCs, large arrow shows Mlh1 focus at base of the XY pairing region. b, Late pachytene spermatocyte. The large arrow identifies an X and Y bivalent without an Mlh1 focus although the number of Mlh1 foci (24) on the autosomal axes at this stage are similar to the number of chiasmata reported for spermatocytes⁴². c, Four bivalents from diplotene oocytes. Mlh1 foci remain at some chiasmata sites (arrow). d, Bivalents from late zyotene/early pachytene oocytes. Animals were from the inbred strain C57Bl/6.

germline of homozygous mutant animals, consistent assure the proper meiotic segregation of the sex chro with a role for Mlh1 in DNA mismatch repair. The elevated levels of microsatellite mutation in Mlh1-deficient mice were similar in degree to the levels previously shown in mice deficient in Pms2, another mismatch repair gene¹⁶. Although the Mlh1-deficient mice are at present too young to draw conclusions on cancer susceptibility, the occurrence of a lymphoma in an Mlh1-deficient animal is consistent with the tumour-prone phenotype reported for Pms2- and Msh2-knockout mice^{16–18}.

Mlh1-deficiency had on both male and female fertility. Mlh1-deficient males produced no spermatozoa, while females produced reduced numbers of oocytes and have thus far proven to be functionally sterile. In contrast, Pms2-deficient mice exhibited male infertility associated with the production of abnormal spermatozoa but no discernible effect on female fertility¹⁶. In addition, the spermatogenesis arrest seen in the Mlh1deficient animals was more uniform than that observed for Pms2 deficiency 16 and suggests a block in first division meiosis.

The abnormal behaviour of homologous chromosomes in Mlh1-deficient mice is consistent with reduced levels of chiasmata. Synapsis appeared to proceed normally, in contrast with meiosis in Pms2-deficient mice¹⁶. However, desynapsis at diplonema in Mlh1-deficient spermatocytes resulted in complete and premature separation of both the XY pair and the autosomal bivalents. By metaphase I, only univalents were observed suggesting either a failure of homologues to cross over or to maintain chiasmata. In contrast, although we do observe univalents in metaphase Pms2-deficient spermatocytes, the degree of univalency is much less than for Mlh1-deficiency. The pairing abnormalities associated with Pms2-deficiency may be sufficient to account for this degree of univalency. Therefore, in the mouse, Mlh1 appears to be required for normal levels of chiasma formation or stabilization, while Pms2 appears to be required for normal homologue pairing.

In parallel studies, we localized Mlh1 in spermatocytes and oocytes using immunostaining. Four aspects of the localization and timing of Mlh1 provide evidence that the meiotic phenotype of Mlh1deficiency reflects a direct role of Mlh1 in crossing over. First, despite variation among mouse strains, the total number of Mlh1 foci on the SCs in midpachynema in both males and females corresponds to the number of previously reported chiasmata⁴² Further, in meiotic analysis of the mouse, a greater number of chiasmata per bivalent were observed in females (in two strains of mice: 1.43 and 1.34 for females; 1.25 and 1.09 for males⁴². Consistent with these studies we observed a comparable difference in the number of Mlh1 foci during mid-pachynema between oocytes and spermatocytes. Second, the timing of stabilization of the number of Mlh1 foci during mid-pachynema in both male and female mice is also consistent with the apparent time of repair DNA synthesis putatively linked to reciprocal recombina-

Third, Mlh1 was localized at the base of the pairing region of the XY where crossing over is necessary to

mosomes in most mammalian species^{26,37-41}. Further, mammalian sex chromosomes proceed through merotic prophase out of synchrony with the autosomes they are the last to synapse and the first to desy napse^{45,46}. The timing of the appearance and disap pearance of an Mlh1 focus on the XY relative to autosomal foci matches this documented asynchrony and is consistent with the premature separation of the X and Y chromosomes in Mlh1-deficient mice.

Fourth, the occurrence of Mlh1 foci at chiasma site A prominent finding was the severe effect that in diplotene oocytes also suggests a direct role for Mlh1 in meiotic crossing over. Although the number of Mlh1 foci at mid-pachynema in oocytes closely corresponds to the previously reported number of chiasmata in diplonema, Mlh1 foci are not present at each chiasma. The absence of Mlh1 at some chiasmata may be due to completion of the Mlh1-dependent step at these locations by the end of pachynema. The disappearance of Mlh1 foci by late pachynema in spermatocytes is also consistent with this interpretation. Pachynema in spermatocytes is prolonged (6 days vs. 3 days in oocytes)47-50 therefore allowing three extra days for completion of Mlh1-dependent processes before the end of pachynema.

> The apparent involvement of Mlh1 in crossing over suggests that Mlh1 may be a component of late RNs, which have been linked to recombination events in mid-to-late pachynema in a variety of organisms^{33,51}. However, mammalian studies have consistently reported a deficiency of late RNs relative to the number of chiasmata in late pachytene nuclei, especially in spermatocytes³⁴. The comparable drop in RNs³⁴ and Mlh1 foci between mid and late pachynema in spermatocytes strengthens the supposition that Mlh1 is a component of late RNs. However, the nature of Mlh1 function in reciprocal recombination and the basis for the apparent localization at crossover sites is not clear. Mlh1 may play a role in the formation, stabilization and/or resolution of Holliday-structure intermediates necessary for crossing over⁵². Alternatively, Mlh1 may be an integral part of a multifunctional complex of proteins involved in recombination between homologous chromosomes. Disruption of Mlh1 could destabilize such a complex, affecting functions in which Mlh1 does not directly participate.

> Although we have focused on the clear connections between Mlh1 and crossing over, we note that the number of Mlh1 foci in early pachytene oocytes exceeds the number of expected crossover events. We speculate that these early pachytene Mlh1 foci are associated with recombination intermediates, only some of which ultimately lead to crossing over. Our results do not provide information on the nature of the recombination intermediate that results in localization of Mlh1 to sites of crossing over. However, we suggest that Mlh1 localization does not simply reflect the presence of DNA mispairs, because the use of inbred mice should minimize the potential for such mismatches. In addition, no significant increase in Mlh1 foci was noted in stermatocytes of mice defi-cient for *Pms2* (unpublished observations), where the associated germline mutator phenotype¹⁶ should have increased the potential for mismatches between homologues. This latter observation also suggests that the

presence of Mlh1 foci is not dependent on the *Pms2* GCTGAGGC; MLH1-T5, GA

Our results show that Mlh1 in the mouse is involved in both DNA mismatch repair and meiotic crossing over. Whereas our studies show that meiosis in Mlh1deficient spermatocytes is characterized by premature separation of homologues, apparently due to unsuccessful completion of recombination and chiasma formation, a previous study showed that Pms2 deficiency was associated with synaptic abnormalities 16. Therefore, the findings reported here and elsewhere 6,9,10,16 indicate that both Mlh1 and Pms2 have roles during DNA mismatch repair and meiosis. However, the different meiotic phenotypes of Pms2 and Mlh1 mice suggest that these two MutL homologues do not fulfill their meiotic functions in concert. Finally, the fertility of Msh2-deficient mice raises the possibility that another MutS-like protein, such as homologues of yeast MSH4 or MSH5 (refs 29,30), act in conjunction with Mlh1 and Pms 2 during meiosis. Alternatively, the two MutL homologues might perform their meiotic functions independent of a MutS homologue.

Methods

Isolation of mouse Mlh1 cDNA. As previously described⁶, a 212-bp fragment from the human Mlh1 was amplified by PCR from human cDNA using degenerate oligonucleotide primers. This human Mlh1 fragment was used to isolate cDNA clones from a mouse teratocarcinoma cDNA library (λ Zap, Stratagene). Dideoxy-sequencing was performed according to the manufacturer (United States Biochemical).

Construction of Mlh1 targeting vector. Using mouse Mlh1 cDNA, a genomic clone was isolated from a 129/Sv strain library (\(\lambda FIXII\), Stratagene) that contained the first five exons of the open reading frame. The genomic insert was subcloned with Notl ends into pBluescript II (SK-) (Stratagene). A 4.7-kb BglII/NotI fragment from the 3' end of the insert was further subcloned into BamHI/NotI digested pBluescript. Following digestion with Sall and blunt ending with T4 polymerase, this 3' homology arm for the Mlh1 targeting vector was cloned into BamHI site (blunted with T4 polymerase), between the HPRT minigene and thymidine kinase gene⁵³. The 5' homology arm, a 4.5-kb EcoRI/XbaI fragment, was subcloned into pBluescript. To prepare the insert for cloning into a Sall site 5' of the HPRT marker, the clone was first digested with Xbal, blunted with T4 polymerase and Sall linkers were ligated. Following digestion with Sall the 5' arm was ligated into the Sall site to generate the Mlh1 targeting vector.

Targeting of Mlh1 in mouse ES cells. AB2.2 ES cells were electroporated with Notl linearized targeting vector and cultured in HAT medium with FIAU selection⁵³. Homologous integration of the targeting construct should introduce an unique Sall restriction site into the Mlh1 locus. Hybridization analysis⁵⁴ of 200 colonies, following digestion with Spel and Sall restriction endonucleases, identified clones which exhibited proper targeting at 5' and 3' ends (Fig. 2).

Generation of Mlh1-deficient mice. Blastocyst microinjection, reimplantation, and the generation of chimaeric males with a high (50–90%) contribution of the ES cell-derived (129Sv/Ev strain) Agouti pigmentation to the coat colour was performed as described^{55,56}. Southern blot analysis was used to identify Mlh1+/- animals among black agouti offspring of chimaeric males, Mlh1-heterozygous siblings were mated to generate Mlh1-/- animals. Subsequent genotyping was performed by PCR using oligonucleotide primers that distinguish the 3' boundary of endogenous and targeted alleles: MLH1-a, AGGAGCTGAT-

GCTGAGGC; MLH1-U, TTTCATCTTGTCACCCGATG; MLH1-T5, GATCTCGACGGTATCGATAAGC. Primers MLH1-a and MLH1-U give a product diagnostic of a untargeted allele that is 258 bp in length, compared to MLH1-a and MLH1-T5, which yield the targeted allele product of 198 bp. PCR was performed in a 25 µl reaction containing 100 ng of genomic DNA and 20 pmol of MLH1-T5 or MLH1-U and 40 pmol of MLH1-a, 0.2 mM of each dNTP, 1.5 mM MgCl₂ and 0.25 U of *Taq Polymerase. Cycling conditions were 94 °C for 4 min, followed by 30 Cycles of 94 °C 1 min, 54 °C for 1 min, 72 °C 1 min. Extension in the last cycle was 72 °C for 3 min.

Antibodies and immunodetection. Antibody production and tests for specificity of Rad51 has been described⁵⁷. The monoclonal antibody to MLH1 was produced by immunizing mice with a recombinant human Mlh1 fusion protein. Hybridoma clones were produced as described^{58,59} and the antibodies purified from supernatants by protein G affinity chromatography. The immunochemical detection procedure was a modification³⁶ of the protocol of Moens et al.⁶⁰. The preparations were incubated overnight at 4 °C with a monoclonal antibody (clone G168-15.3) against human MLH1 (diluted 1:400) and a polyclonal antibody against human RAD51 raised in rabbit⁵⁷ (diluted 1:200). The MLH1 antibody was detected with goatanti-mouse-IgG conjugated with rhodamine (Pierce) and the RAD51 antibody with goat-anti-rabbit-IgG conjugated with fluorescein isothiocyanate (FITC, Sigma). All preparations were counter stained with 4',6-diaminino-2-phenylindole (DAPI). Preparations were examined and digitally imaged as described36.

Western blot analysis. Culture of cells isolated from 15-day embryos is described elsewhere¹⁶. For Western blot analysis, protein from cultured cell extracts was resolved by SDS-PAGE, electro-blotted to Immobilion P transfer membrane (Millipore), and treated with Anti-MLH1 antibodies. Bound MLH1 antibody was detected by chemiluminescence (Renaissance, DuPont) using goat anti mouse IgG horseradish peroxidase conjugate (Pierce).

Histology of testis and ovary. Preparation of testis for histological analysis is described elsewhere 16 . All surrounding tissue was removed from the ovaries prior to overnight fixation in Zincbuffered formalin. The fixed samples were embedded in paraffin wax, sectioned (4 μ m), mounted and stained with either hematoxylin and esoin, or periodic acid, Schiff's and hematoxylin.

Single molecule PCR analysis. The PCR primers used for D9Mit67 were as described16. Primers for amplification of locus D1Mit355 in the first round were P1(5'-TGAAAAGAC-CTTTTCTCAAATAGTG-3') and P2(5'-CTTTGATTCT-GAAATATACAGCAA-3'). In the second round P2 was replaced with P3(5'-TAGGAACTGTTTTGTTGTTTTACAC-3'). The PCR buffer was described previously35. Volumes for first round reactions were 20 µl. From the first round 2 µl of product was added to 40 µl of PCR buffer for the second round. After an initial denaturation at 94 °C for 4 min, cycling conditions were for the first round of PCR an annealing step of 60 °C for 1 min (57 °C for D1Mit355), an extension step of 72 °C for 2 min and a denaturation step of 94 °C for 30 s (30 cycles). The second round of 25 cycles was identical for both loci and annealing and extension was carried out at 60 °C for 1.5 min. The last cycle of each round was an extension at 72 °C for 7 min.

Animals for Mlh1 localization. All normal mice were from the C57BL/6 inbred strain. Over 100 nuclei were imaged from 4 juvenile (18–21 days old) and 2 adult males, and over 150 nuclei were imaged from oocytes from 16–19 day old fetuses from 8 different litters. In addition, 50 nuclei were examined from an Mlh1-deficient mouse from a mixed 129/C57 genetic background. Surface spreads were prepared as described³⁶.

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